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1 **Calcium isotopes as a record of the marine calcium cycle versus carbonate diagenesis**
2 **during the late Ediacaran**

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17

18 **Highlights**

- 19 • $\delta^{44}\text{Ca}$ from limestones aged ~550–539 Ma that host the earliest skeletal animal
20 fossils
- 21 • Negative shift in $\delta^{44}\text{Ca}$ that lasted at least 11–14 Myrs
- 22 • Unlikely to record a transition towards more sediment-buffered carbonate
23 diagenesis

- May record enhanced continental weathering or evaporite deposition
- May record a change in timing of dolomitisation

Keywords: Calcium isotopes; carbonate rocks; marine calcium cycle; analytical methods; Ediacaran; biomineralisation

Abstract

Calcium isotope ratios in ancient carbonate rocks can provide insight into the global marine calcium cycle as well as local conditions during carbonate mineral precipitation and diagenesis. We compare two extraction techniques for the separation of calcium from other ions before $\delta^{44}\text{Ca}$ analysis, using an automated ion chromatograph and using manual gravity columns. The two techniques produce the same $\delta^{44}\text{Ca}$ within error (2σ). We present 31 $\delta^{44}\text{Ca}$ analyses of carbonate rocks from the Nama Group, Namibia, which record a negative shift in $\delta^{44}\text{Ca}$ of 0.35‰ between ~ 550 and ~ 547 Ma, from -1.25‰ to -1.60‰ , followed by persistently low $\delta^{44}\text{Ca}$ ($-1.48 \pm 0.06\text{‰}$) between ~ 547 and 539 Ma. Very low $\delta^{44}\text{Ca}$ ($< -1.5\text{‰}$) are commonly interpreted to represent the preservation of local aragonite that has recrystallized to calcite under sediment-buffered conditions (where the composition of the diagenetic carbonate product is determined mainly by the original sediments). The shift in $\delta^{44}\text{Ca}$ across the Nama Group could therefore represent a change from fluid-buffered diagenesis (where the composition of the diagenetic carbonate mineral is determined mainly by the fluid) to sediment-buffered diagenesis. However, this interpretation is not consistent with either potential geochemical indicators of diagenesis (e.g., $\delta^{18}\text{O}$), or changes in large-scale fluid-flow as predicted from sequence stratigraphy.

47 We consider alternative interpretations for generating changes in the $\delta^{44}\text{Ca}$ of ancient
48 carbonate rocks including enhanced continental weathering, increases in evaporite
49 deposition, and changes in the style of dolomitisation.

50

51 **1. Introduction**

52 Calcium isotope ratios (the ratio of $^{44}\text{Ca}/^{40}\text{Ca}$ and reported versus a standard as
53 $\delta^{44}\text{Ca}$) measured in ancient carbonate minerals and rocks such as limestone and dolomite,
54 offer a potential tool to investigate the workings of the ancient marine calcium cycle as well
55 as to explore carbonate mineral deposition and subsequent diagenesis (Blättler et al., 2012,
56 2011; Bradbury and Turchyn, 2018; Higgins et al., 2018; Husson et al., 2015). The calcium
57 isotope ratio of seawater, $\delta^{44}\text{Ca}_{\text{sw}}$, is controlled by the balance of fluxes of calcium into and
58 out of seawater, and the calcium isotopic composition of those fluxes. The main source of
59 calcium to seawater is riverine input, although hydrothermal fluids may account for ~17%
60 of total calcium input to the ocean (Tipper et al., 2010). The main output flux is the
61 deposition of carbonate-bearing sediments, but evaporite deposition and alteration of
62 oceanic crust are also important sinks for calcium.

63 Calcium isotopes preserved in carbonate rocks reflect both the $\delta^{44}\text{Ca}$ of the seawater
64 or fluid from which carbonate minerals precipitated, as well as the kinetic calcium isotope
65 fractionation during precipitation ($\Delta^{44}\text{Ca}_{\text{local}}$), which is determined by the carbonate
66 mineral precipitating (e.g., calcite, aragonite or dolomite) and its rate of precipitation
67 (Blättler et al., 2012; Tang et al., 2008). At steady state, the dominant control on the
68 calcium isotope composition of seawater ($\delta^{44}\text{Ca}_{\text{sw}}$) is the average $\delta^{44}\text{Ca}$ of global buried
69 carbonate minerals ($\delta^{44}\text{Ca}_{\text{global}}$). However, carbonate rocks are also the major archive used

70 to reconstruct seawater $\delta^{44}\text{Ca}$ in deep time. Therefore, to interpret $\delta^{44}\text{Ca}_{\text{sw}}$ from $\delta^{44}\text{Ca}$ in
71 carbonate rocks, a *local* calcium isotope fractionation factor must be assumed based on the
72 inferred original (primary) and dominant mineralogy. Although early $\delta^{44}\text{Ca}$ data were
73 commonly interpreted to reflect changes in the global calcium cycle, i.e., the relative
74 sources and sinks of calcium to the ocean (Blättler et al., 2012, 2011; Hinojosa et al., 2012;
75 Payne et al., 2010), more recently, $\delta^{44}\text{Ca}$ data have been interpreted to record calcium
76 isotope signatures acquired during diagenesis at the site of sediment lithification (Ahm et
77 al., 2019, 2018; Higgins et al., 2018; Husson et al., 2015).

78 The late Ediacaran was a time of dramatic environmental and ecological change,
79 including oscillating redox conditions, extensive evaporite deposition, and the appearance
80 of skeletal animals (Fike and Grotzinger, 2008; Tostevin et al., 2019; Wood, 2011; Wood et
81 al., 2017b). We present $\delta^{44}\text{Ca}$ data from the Nama Group, Namibia, that record a negative
82 shift in $\delta^{44}\text{Ca}$ of $\sim 0.35\text{‰}$ between ~ 550 and ~ 547 Ma, followed by persistently low $\delta^{44}\text{Ca}$
83 ($-1.48 \pm 0.06\text{‰}$) from ~ 547 to ~ 539 Ma. We combine these data with other major element
84 and isotopic geochemical data from the same samples to evaluate several potential
85 explanations for this $\delta^{44}\text{Ca}$ shift, and discuss the application of calcium isotope systematics
86 in deep time.

2. Geological and geochemical context

2.1 Sequence stratigraphy and diagenetic history of the Nama Group

The Nama Group was deposited in two inter-connected sub-basins, the southern Witputs and the northern Zaris, separated by a zone of depositional thinning representing the “Osis Arch” paleo-bathymetric high (Germs, 1974). Unrestricted connection to the open ocean has been suggested because of the preservation of $\delta^{13}\text{C}$ excursions in time-equivalent sections, and normal marine rare earth element signatures (Bowyer et al., 2017; Kaufman et al., 1991; Tostevin et al., 2016b; Wood et al., 2015). The Zaris Formation (Kuibus Subgroup) in the Zaris sub-basin was sampled at Zebra River Farm, and the Urusis Formation (Schwarzrand Subgroup) in the Witputs sub-basin was sampled at Swartkloofberg Farm and at Swartpunt Farm. The base of the Nama Group is diachronous, but has been estimated to have been deposited between 553 and 550 Ma (Ries et al., 2009; Saylor et al., 1998). An ash bed in the Hoogland Member yields a depositional age of 547.32 ± 0.65 Ma (Bowring et al., 2007; Grotzinger et al., 1995). Therefore, deposition of the Omkyk and Hoogland members of the Zaris Formation likely spans 3–6 Myrs. Ash beds at Swartpunt Farm yield ages of 540.095 ± 0.099 Ma and 538.99 ± 0.21 Ma for the top of the Schwarzrand Subgroup (Linnemann et al., 2019). Overall, the Nama Group therefore spans 11–14 Myrs.

The Nama Group hosts terminal Ediacaran skeletal fauna, *Cloudina*, *Namacalathus* and *Namapoikia*, as well as soft-bodied Ediacaran fossils and increasing trace fossil evidence for motility towards the top of the section (Germs, 1972; Grant, 1990; Grotzinger et al., 2000; Jensen et al., 2000; Wood et al., 2002). At the deep inner-ramp locality, Zebra River, the Lower Omkyk Member is dominated by grainstones. In the transgressive systems

110 tract of the Upper Omkyk Member, thrombolite-stromatolite reefs nucleate, forming
111 laterally continuous biostrome layers (Grotzinger et al., 2000). *Cloudina* and *Namacalathus*
112 can be found within thrombolite heads and lag beds within inter-reef shales. Towards the
113 top of the Upper Omkyk Member the section shallows into grainstone-dominated facies
114 with subordinate shale horizons, containing thinner, discontinuous biostrome microbial
115 reef systems, and some large *Namacalathus* <35 mm (Penny et al., 2016). The Hoogland
116 Member contains storm dominated laminites and heterolithics, shallowing towards
117 grainstone-dominated facies.

118 The Pinnacle Reefs at Swartkloofberg Farm were deposited in a mid-ramp setting
119 within a transgressive systems tract, initiated on the flooded surface of the Huns Platform.
120 After termination of reef growth, the reefs were enveloped by shales, and together these
121 facies form the Feldschuhhorn Member. The Pinnacle Reefs host communities of skeletal
122 animals of varied sizes, including aggregations of *Namacalathus* up to 12 mm in diameter
123 (Wood et al., 2015).

124 Swartpunt Farm, which encompasses the Spitzkopf Member of the Schwarzrand
125 Subgroup, transitions from low-energy outer-ramp setting at the base, towards an inner-
126 ramp environment, followed by a deepening to outer ramp conditions, and a transition
127 towards a mid- to inner-ramp setting at the top of the section. Thin-bedded calcisiltite beds
128 at the base of Swartpunt host *Namacalathus*, large *Cloudina*, and a diversity of carbonate
129 microbialites, and are interpreted to have been deposited in a low-energy deeper ramp
130 setting (Narbonne et al., 1997; Saylor, 2003). The limestone is overlain by thick beds of
131 green mudstone and coarse sandstone, deposited in a deltaic environment, containing
132 burrows and soft-bodied fossils including *Swartpuntia* and *Pteridinium* (Narbonne et al.,

1997). Overlying this there is a 5–10 m interval of thinly bedded siltstone, sandstone and limestone with ripple cross-lamination, deposited above fair weather wave base in an inner-ramp setting. Overlying shales may represent deepening to an outer-ramp environment. These give way to limestones and dolomites with dm-scale thrombolites, deposited at or below storm wave base in a mid-ramp setting (Jensen and Runnegar, 2005). Towards the top of Swartpunt is a highstand system tract containing flaggy, laminated limestones with small (<5 mm) *Cloudina riemkeae* and thrombolites.

The paragenetic sequence of carbonate rocks from the Nama Group has been described in detail from the *Cloudina* reefs at Driedoornvlagte Farm, coeval in part with the Zebra River section, identifying six successive cement generations (Wood et al., 2018). *Cloudina* are commonly preserved as neomorphosed calcite, with micro-dolomite inclusions, and in some cases are completely dolomitised. Large, acicular cements form botryoids and occlude pore space. These commonly nucleate on skeletal fossils, or intergrow with geopetal sediment, and are interpreted to represent an early marine precipitate of originally aragonitic mineralogy (Grant, 1990). This is followed by a thin, isopachous dolomite cement. This is post-dated by a cloudy, inclusion-rich low-Mg calcite cement, interpreted to have precipitated in pore fluids that were in open contact with seawater. Remaining pore space is occluded by a clear, length-slow, blocky low-Mg, calcite, typical of low-Mg pore waters, suggesting a burial origin. Early marine cements are commonly interpreted to be aragonitic, and later neomorphosed to calcite (Grant, 1990; Grotzinger and James, 2000; Wood et al., 2018).

2.2 Geochemical background of the Nama Group samples

The Nama Group was deposited coincident with the final recovery from the Shuram-Wonoka anomaly, an enigmatic global $\delta^{13}\text{C}$ excursion famed for reaching unusually low values (around -12‰). The Shuram-Wonoka anomaly has been variably interpreted to result from either: 1) the oxidation of a large pool or reduced carbon such as methane hydrates or dissolved organic carbon (Bjerrum and Canfield, 2011; Husson et al., 2015; Rothman et al., 2003), 2) a global increase in the burial of authigenic carbonate (Cui et al., 2017; Schrag et al., 2013), or 3) globally synchronous changes in burial or meteoric diagenesis (Derry, 2010; Knauth and Kennedy, 2009). Some sections in the Nama Group capture the tail end of the Shuram excursion (e.g., at Brak and Grens; Wood et al., 2015), but the Nama Group at Zebra River captures the post-excursion return to positive $\delta^{13}\text{C}$. In the Schwarzsand Subgroup, $\delta^{13}\text{C}$ remains stable around 1‰ .

Iron speciation and cerium anomaly measurements have previously been used in this section as proxies for local water column redox conditions (Tostevin et al., 2016b; Wood et al., 2015). These data indicate that Zebra River was predominantly well-oxygenated, with temporary anoxic-ferruginous periods (Wood et al., 2015), which may reflect the development of sluggish circulation or upwelling of anoxic deeper waters associated with marine transgression (Bowyer et al., 2017). Additionally, positive cerium anomalies suggest that some depositional intervals were oxygen-poor and manganoan (Tostevin et al., 2016). The Pinnacle Reefs experienced persistent well-oxygenated conditions (Tostevin et al., 2016b; Wood et al., 2015). Swartpunt was largely oxygenated, with anoxic ferruginous conditions restricted to two highstand carbonate horizons. Sediments containing independent evidence for deposition under an oxygenated water

column were selected for this study, and all samples have low total organic carbon (TOC) of <0.2 wt.% (Wood et al., 2015).

$\delta^{238}\text{U}$ data from carbonate rocks at Zebra River transition from modern-like seawater values (-0.4‰) in the Lower Omkyk Member, to very negative values in the Upper Omkyk and Hoogland Members (-0.81‰) (Tostevin et al., 2019). This has been interpreted and modelled to represent a transition towards globally widespread anoxic bottom waters that covered at least a third of the global sea floor. Further, this $\delta^{238}\text{U}$ transition appears to be recorded globally in sediments of the same age (Zhang et al., 2018), supporting preservation of a primary global seawater $\delta^{238}\text{U}$ signal. Existing sulfur isotope data from carbonate associated sulfate at Zebra River show a general increasing trend in $\delta^{34}\text{S}_{\text{CAS}}$ from the Omkyk to the Hoogland Members (Tostevin et al., 2017). This correlates with contemporaneous sections from other basins, suggesting a global change in the sulfur cycle that drove an increase in global marine $\delta^{34}\text{S}$ (Cui et al., 2016b, 2016a; Fike and Grotzinger, 2008). This has been interpreted to reflect a change in weathering fluxes or sources, and/or an increase in the global pyrite burial flux.

3. Methods

Hand samples were collected at one to five meter intervals along with stratigraphic logs noting lithology and paleoecology. Weathered edges were removed and samples were sawed in half to reveal a fresh surface. Powders from Zebra River for $\delta^{44}\text{Ca}$ analysis were drilled with a dremel microdrill, avoiding visually recrystallised areas. Bulk-rock powders were analysed from the Pinnacle Reefs and Swartpunt sections.

Major element concentrations (Sr, Ca, Mg and Mn) in the carbonate portion of the sample were determined using sequential leaching in 2% HNO_3 , and analysed via Inductively coupled plasma optical emission spectrometer (ICP-OES) at the Cross-Faculty Elemental Analysis Facility, University College London. The sequential leaching method involves pre-leaching 20% of the sample, followed by a 40% leach that is retained for analysis (method is described in full in Tostevin et al., 2016a). These major element data are available for Zebra River section as well as four other localities in the Nama Group (the Kuibis Subgroup at Brak, Omkyk and Zwartmodder, and the Schwarzrand Subgroup at Swartpunt; Wood et al., 2015). Carbon isotopes have been previously reported in Wood et al., (2015).

These data were used to inform sample selection for calcium isotope analysis. Samples with a wide range of Sr contents, including those that were anomalous compared with adjacent samples, were selected. Selected samples included a range of facies and textures, including microbialites, grainstones and laminites. Dolomitised or impure samples (low $\%\text{CaCO}_3$) were avoided. Calcium isotope ($^{44}\text{Ca}/^{40}\text{Ca}$) analysis was conducted at the University of Cambridge using a Thermo Scientific Triton Plus MC-Thermal Ionisation Mass Spectrometer. Sample aliquots containing six μg of calcium were combined

216 with a ^{42}Ca - ^{48}Ca double-spike at a ratio of 10:1 (sample-to-spike) in acid-cleaned Teflon
217 vials. The 48:42 ratio of the double-spike is 1:1, similar to the optimum ratio of 3:2 for a
218 ^{42}Ca - ^{48}Ca double-spike (Rudge et al., 2009). Solid samples were dissolved in dilute ultra-
219 pure acetic acid for 1 h, before being converted to nitrates and then combined with the
220 double spike. The samples were then dried and re-dissolved in 0.5% nitric acid and calcium
221 was separated using either a Dionex ICS 5000+ HPIC coupled with a Dionex AS-AP fraction
222 collector or a gravity column (Bio-Rad AG50W-X8) setup for method validation.

223 After the separation of the calcium using either the Dionex ICS 5000+ HPIC or the
224 gravity columns, 4 μg of calcium was loaded on an outgassed 0.7 mm Rhenium filament
225 with 0.5 μl of 10% trace metal purity Phosphoric acid as an activator. The samples are run
226 using the analytical method previously described in Bradbury and Turchyn (2018). The
227 average external 2σ standard deviation over nine months on the standard NIST915B was
228 0.10‰ ($n = 82$). All $\delta^{44}\text{Ca}$ discussed in the text are reported as ratios of ^{44}Ca to ^{40}Ca relative
229 to modern seawater, but in Table 1 the data are also reported relative to other commonly
230 used calcium standards (BSE and 915a).

231

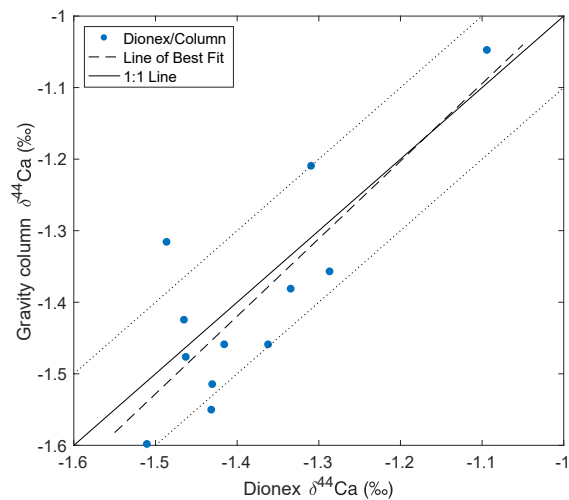
232 **3.1 Method Validation**

233 Each sample was run for its calcium isotopic composition four times, using two
234 extraction techniques; separating the calcium using an automated ion chromatograph and
235 separating the calcium using manual gravity columns. The samples run using the Thermo
236 Scientific Dionex ICS 5000+ HPIC were run through a high-capacity carboxylate-
237 functionalised column (Dionex CS-16) using 30 mM methyl-sulfonic acid (MSA) at a flow

238 rate of 1 ml/min. The conductivity of the samples was continuously measured during the
239 separation, and a minimum peak slope of 0.003 $\mu\text{S/s}$ determined the sample collection
240 during a set time window. The method is similar to the published work of Schmitt et al.,
241 (2009), and has been published in Bradbury et al., (2018) and Bradbury and Turchyn,
242 (2018).

243 These analyses were compared to samples that were separated through a
244 traditional gravity column setup using Bio-Rad AG50W-X8 resin. The gravity columns
245 were created from cut down 5 ml pipettes with a reservoir size of 2 ml and with a 20 μm PE
246 frit installed. They were filled to just below 1 ml with slurried AG50W-X8 resin. The
247 columns were pre-cleaned using 2 ml of 4 M HCl, 6 M HNO_3 , 4 M HCl and water, before
248 being preconditioned with 1.5 M HNO_3 . The columns had a flow rate of approximately 1
249 ml/hour. The sample was loaded and then eluted with 1.5 M HNO_3 . The columns were
250 calibrated by measuring the concentration of the ions eluting off the column using the
251 Dionex ICS 5000+. The calibration was setup to collect 100% of the calcium and maximise
252 the separation of calcium from magnesium and strontium, whilst at the same time
253 minimising the time taken to complete the column chemistry. Initially a series of twelve
254 carbonate powders from the Nama Group were dissolved in 5% ultrapure acetic acid. An
255 aliquot containing 12 μg of calcium was then spiked at a ratio of 10:1 (sample to spike)
256 with the double spike and dried. The dried sample was re-dissolved in 1.5 M HNO_3 and
257 separated using the calibrated gravity columns. The 4 μg of the collected calcium was
258 loaded per filament either singularly ($n=6$), in duplicate ($n=6$), or triplicate ($n=12$). The
259 measured sample data is shown graphically in Figure 1, where the $\delta^{44}\text{Ca}$ from the Dionex
260 ICS 5000+ HPIC and the column separations are compared.

261



262

263 **Figure 1:** Cross plots of the twelve samples run on the Dionex ICS 5000+ and gravity
264 columns (n=12, R²=0.65, p-value = 0.0119). The dotted lines represent the long-term 2σ of
265 the measurement of calcium isotopes. **A t-test comparison of the slopes of the measured**
266 **data and the 1:1 line shows no statistically significant difference.**

267

268 **4. Results**

269 We report 31 new $\delta^{44}\text{Ca}$ analyses from the Nama Group, which range from -1.60‰
270 to -1.07‰ . At Zebra River, $\delta^{44}\text{Ca}$ progressively decreases from the base of the section
271 through the Kuibis Subgroup from -1.25‰ to -1.60‰ , an overall decrease of 0.35‰
272 (Figure 2). There is one outlying data point, with a particularly high $\delta^{44}\text{Ca}$ of -1.09‰ (at
273 106 m). At the Pinnacle Reefs and Swartpunt, $\delta^{44}\text{Ca}$ remains low through the Schwarzrand
274 Subgroup, between -1.39‰ and -1.59‰ (average = -1.48 ± 0.06). Two partially

275 dolomitised samples have a $\delta^{44}\text{Ca}$ that is $\sim 0.4\text{‰}$ higher than the surrounding samples
276 ($\sim -1.08\text{‰}$ at 5 m and $\sim -1.07\text{‰}$ at 10 m), and these have been excluded from consideration
277 due to the known positive $\delta^{44}\text{Ca}$ offset associated with dolomitisation (Fantle and DePaolo,
278 2007; Turchyn and DePaolo, 2011). The samples are all laminite heterolithics, grainstones,
279 microbialites, or recrystallized limestones, and changes in $\delta^{44}\text{Ca}$ do not appear to correlate
280 with changes in these lithologies (Figure 2).

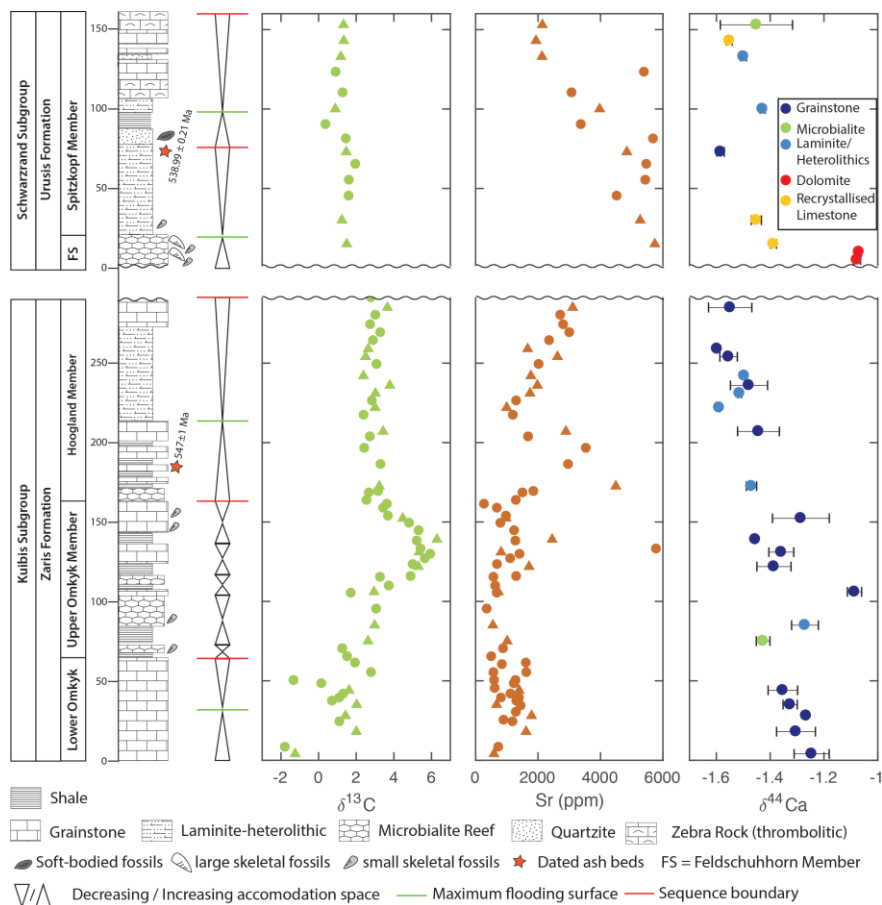


Figure 2: From left to right: Stratigraphic log, sequence stratigraphy, $\delta^{13}\text{C}$ data, Sr content and $\delta^{44}\text{Ca}$ for the Nama Group. Triangle symbols for $\delta^{13}\text{C}$ data and Sr content highlight samples for which there is also $\delta^{44}\text{Ca}$ data. The $\delta^{44}\text{Ca}$ data are colour coded according to the facies of individual samples. The Kuibis Subgroup was sampled at Zebra River, the Feldschuhhorn Member (FS) was sampled at the Pinnacle Reefs, and the Spitzkopf Formation was sampled at Swartpunt Farm. Ash bed dates from Hoogland Member and Spitzkopf Member are from Bowring et al., (2007) and Linnemann et al., (2019). Sequence stratigraphy is from Adams et al., (2005), Saylor (2003) and Saylor et al., (1998). **The wavy line in the stratigraphic column does not indicate a substantial erosional unconformity, but an absence of samples from the intervening stratigraphy.**

Strontium content in the carbonate rocks is consistent throughout the Omkyk Member, with an average of 1000 ppm, but rises through the Hoogland Member reaching a peak of ~4000 ppm. In the Schwarzrand Subgroup, the strontium content declines from a high of ~6000 ppm to ~2000 ppm close to the Ediacaran-Cambrian boundary. This increase in strontium content is consistent across multiple sections of the Nama Group (Wood et al., 2015; Figure 3). At Zebra River, scatter overprints the overall trend in Sr content, with some exceptionally high values in the middle of the section. $\delta^{44}\text{Ca}$ correlates with Sr content ($R^2=0.27$), $\delta^{34}\text{S}_{\text{CAS}}$ ($R^2=0.53$) and $\delta^{18}\text{O}$ ($R^2=0.40$) measured on the same samples at the $P<0.05$ level, but with significant scatter around the trend (Figure 4b, 4f and 4k). In contrast, there is no significant correlation between $\delta^{44}\text{Ca}$ and other geochemical data ($P>0.05$), including Mn content, $\delta^{13}\text{C}$, TOC and Mg/Ca (Figure 4a and 4c-e).

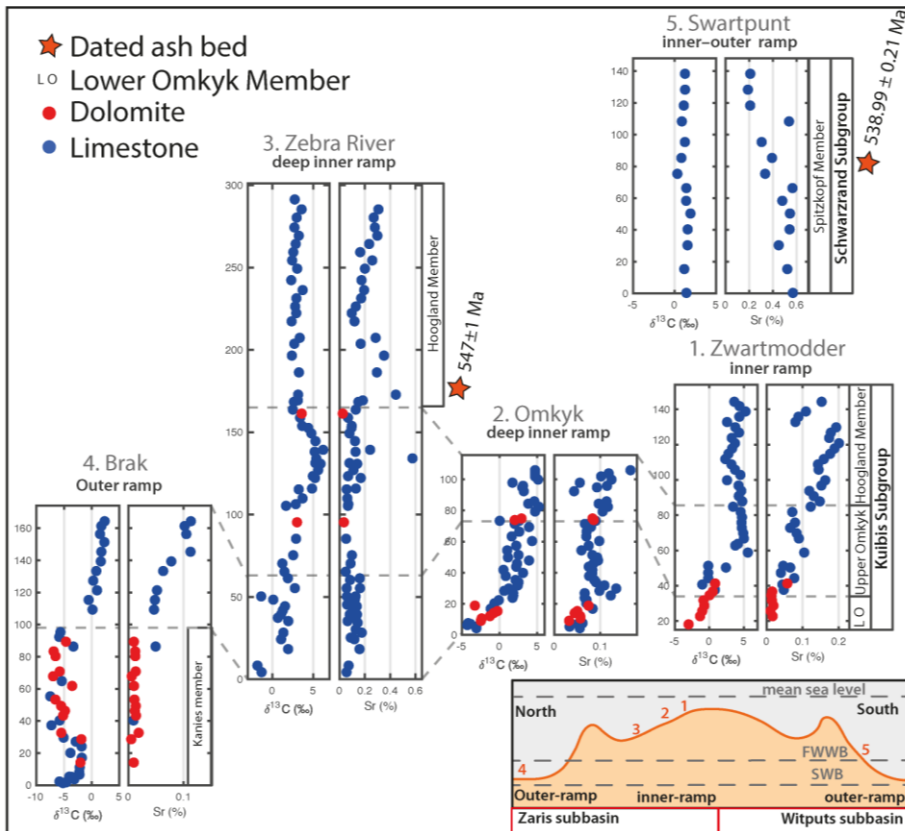


Figure 3: Sr concentrations for four sections from the Kuibis Subgroup, and one section from the Schwarzsrand Subgroup. Carbon isotopes from Wood et al., (2015). Ash bed dates from Hoogland Member and Spitzkopf Member are from Bowring et al., (2007) and Linnemann et al., (2019).

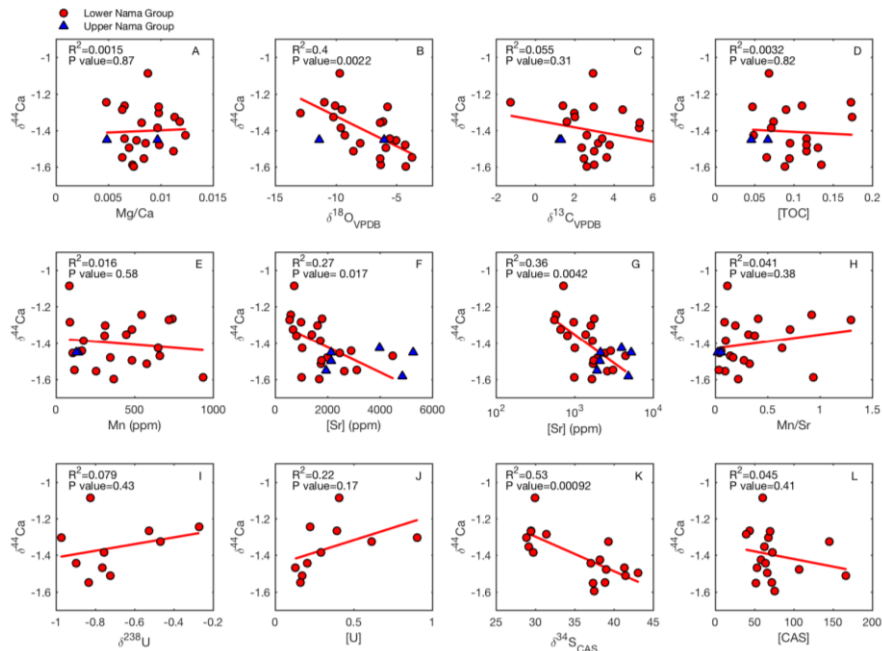


Figure 4: Cross plots of $\delta^{44}\text{Ca}$ against other geochemical parameters measured on the same samples. Red circles are from the Kuibis Subgroup, and blue triangles are from the Schwarstrand Subgroup. P-values and R^2 values for linear best fit lines are noted for the Kuibis Subgroup. Sr content is shown twice, once on a linear scale (F) and once on a log-linear scale (G).

Table 1: $\delta^{44}\text{Ca}$ data for each sample relative to modern seawater (SW), 915a (1.94‰ lower than SW), and BSE (0.94‰ lower than SW), along with the standard deviation (1σ) for the $\delta^{44}\text{Ca}$ data, and the strontium content.

Locality	Sample name	Height (m)	$\delta^{44}\text{Ca}$ (SW)	$\delta^{44}\text{Ca}$ (915a)	$\delta^{44}\text{Ca}$ (BSE)	1σ	Sr content (ppm)
----------	-------------	------------	-----------------------------	-------------------------------	------------------------------	-----------	------------------

<i>Zebra River</i>	LO2	4	-1.25	0.69	-0.31	0.065	590
	LO4	18	-1.30	0.64	-0.36	0.072	1617
	LO6	28	-1.27	0.67	-0.33	0.000	1794
	LO7	35	-1.33	0.61	-0.39	0.026	674
	LO11	44	-1.35	0.59	-0.41	0.055	1396
	OS2-3	75	-1.43	0.51	-0.49	0.025	1014
	OS2-4	85	-1.27	0.67	-0.33	0.050	555
	ZR5	106	-1.09	0.85	-0.15	0.026	730
	ZR9	122	-1.39	0.55	-0.45	0.063	1714
	OS2-9	131	-1.36	0.58	-0.42	0.046	818
	OS2-10	139	-1.46	0.48	-0.52	0.000	2452
	ZR18	152	-1.29	0.65	-0.35	0.105	979
	ZR31	173	-1.47	0.47	-0.53	0.019	4489
	ZR38	207	-1.44	0.50	-0.50	0.078	2892
	UH2	222	-1.59	0.35	-0.65	0.000	998
	UH4	231	-1.51	0.43	-0.57	0.007	1744
	UH5	236	-1.48	0.46	-0.54	0.069	1981
	UH6	242	-1.50	0.44	-0.56	0.000	1772
	UH8	254	-1.55	0.39	-0.61	0.032	2622
	UH9	259	-1.60	0.34	-0.66	0.000	1668
	UH14	285	-1.56	0.38	-0.62	0.000	
<i>Pinnacle Reefs</i>	PR7	5	-1.08	0.86	-0.14	0.013	
	PR	10	-1.07	0.87	-0.13	0.009	
	PR6	15	-1.39	0.55	-0.45	0.011	
<i>Swartpunt</i>	SW2	30	-1.45	0.49	-0.51	0.019	5270
	SW6	73	-1.58	0.36	-0.64	0.014	4845
	SW9	100	-1.43	0.51	-0.49	0.010	3973
	SW12	133	-1.50	0.44	-0.56	0.011	2128
	SW13	143	-1.55	0.39	-0.61	0.011	1936
	SWP15	153	-1.45	0.49	-0.51	0.134	2138

324 5. Discussion

325 The decrease of 0.35‰ in the calcium isotopic composition of carbonate rocks
326 recorded through the Kuibis Subgroup is substantial, and of a similar magnitude to the
327 change in $\delta^{44}\text{Ca}$ through the end-Ordovician mass extinction (0.5–0.6‰; Holmden et al.,
328 2012), Cretaceous anoxic events (0.2–0.4‰; Blättler et al., 2011), and the Permo–Triassic
329 Boundary (0.3‰; Payne et al., 2010). Following the progressive transition towards lower
330 $\delta^{44}\text{Ca}$ across the Kuibis Subgroup, $\delta^{44}\text{Ca}$ remains low (around –1.5‰) throughout the
331 Schwarstrand Subgroup, up to the Ediacaran–Cambrian Boundary.

332 A similar negative shift in $\delta^{44}\text{Ca}$ is recorded in contemporaneous rocks from South
333 China (Sawaki et al., 2013), although those data come from five samples within a mixed
334 dolomite and limestone succession, making it difficult to discern mineralogical controls
335 from trends in seawater calcium isotopic composition. Pruss et al., (2018) report $\delta^{44}\text{Ca}$
336 from bulk rock samples from the Omkyk Member of the Nama Group that overlap with the
337 range of $\delta^{44}\text{Ca}$ in this study (–1.07‰ to –1.59‰), but the $\delta^{44}\text{Ca}$ data come from fossil
338 samples and are not presented stratigraphically. $\delta^{44}\text{Ca}$ data from the Wonoka Formation,
339 deposited during the Shuram–Wonoka carbon isotope excursion, identify a prominent
340 negative excursion, reaching from –0.8‰ to –1.9‰, before recovering to –0.8‰ (Husson
341 et al., 2015). Dolomitised samples at the top of the Wonoka Formation may be
342 contemporaneous with limestone samples at the base of the Nama Group, but the differing
343 mineralogy makes it difficult to directly compare the $\delta^{44}\text{Ca}$ values.

344 Assuming the Nama Group has not experienced diagenetic alteration with fluid that
345 has an exceptionally low $\delta^{44}\text{Ca}$ (Gussone et al., 2005), then the very low $\delta^{44}\text{Ca}$ recorded in
346 the upper Kuibis and Schwarstrand subgroups can be interpreted to result from two

possible **endmember** scenarios: 1) local carbonate deposition associated with a larger local calcium isotopic fractionation ($\Delta^{44}\text{Ca}_{\text{local}}$) of around -1.7‰ , in an ocean with a similar $\delta^{44}\text{Ca}_{\text{sw}}$ to today, or 2) local carbonate deposition associated with a smaller $\Delta^{44}\text{Ca}_{\text{local}}$ of around -1.25‰ , in an ocean with $\delta^{44}\text{Ca}_{\text{sw}}$ that is $0.4\text{--}0.5\text{‰}$ lower than today.

5.1 Scenario one: Preservation of local aragonite $\delta^{44}\text{Ca}$

In an ocean with a similar $\delta^{44}\text{Ca}$ to today, preservation of very low $\delta^{44}\text{Ca}$ ($<-1.5\text{‰}$) in carbonate rocks requires a large $\Delta^{44}\text{Ca}_{\text{local}}$ during carbonate mineral deposition (e.g., scenario 1). A local change from calcite towards aragonite deposition, or an increase in the precipitation rate of carbonate minerals, could drive changes in the Sr content and $\delta^{44}\text{Ca}$ of carbonate rocks in the direction and magnitude recorded across the Kuibis Subgroup (Figure 5a; Farkaš et al., 2016; Tang et al., 2008). It is likely that the original sedimentary mineralogies in the Nama Group were dominantly aragonite, consistent with petrographic work that has identified a dominantly aragonitic primary mineralogy for many, but not all, components including large botryoidal cements **and crystal fans** (Grant, 1990; Grotzinger et al., 2005; Wood et al., 2018).

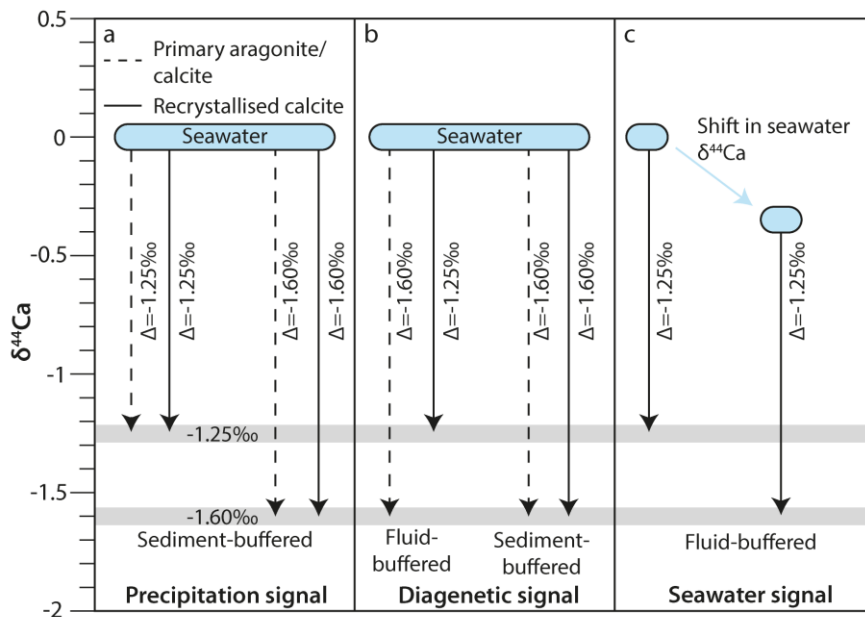


Figure 5: Schematic showing three possible models to explain the shift towards lower $\delta^{44}\text{Ca}$ across the Nama Group. Panel a) shows a change in fractionation during precipitation of primary minerals from seawater, resulting from either a change in mineralogy or precipitation rates, preserved in recrystallized calcite through sediment-buffered diagenesis. Panel b) shows a diagenetic interpretation, where the primary aragonite $\delta^{44}\text{Ca}$ is the same, but differences in $\delta^{44}\text{Ca}$ are introduced during recrystallisation to calcite under fluid- and sediment-buffered diagenesis (see section 5.1). Panel c) illustrates a change in seawater $\delta^{44}\text{Ca}$, which could be preserved in carbonate rocks during either fluid- or sediment-buffered diagenesis (see section 5.2).

375 During recrystallisation of primary aragonite to calcite, the $\delta^{44}\text{Ca}$ of the calcite is
376 determined by the $\delta^{44}\text{Ca}$ of the pore fluid from which it precipitates. Under conditions of
377 high fluid flow, pore fluids can be buffered by seawater Ca (fluid-buffered conditions). In
378 contrast, when pore fluids become isolated from seawater, the $\delta^{44}\text{Ca}$ of the pore fluid can
379 be buffered by the dissolution of the primary aragonite (sediment-buffered conditions)
380 (Higgins et al., 2018). For primary aragonite $\delta^{44}\text{Ca}$ to be preserved it is thought that
381 recrystallisation of the primary aragonite to calcite must occur under sediment-buffered
382 conditions, such that the buried calcite should retain its primary aragonite $\delta^{44}\text{Ca}$. If instead,
383 recrystallisation occurs under fluid-buffered conditions, the buried calcite can acquire a
384 new $\delta^{44}\text{Ca}$ that reflects the smaller $\Delta^{44}\text{Ca}_{\text{local}}$ between newly precipitated calcite and the
385 pore fluid (Ahm et al., 2018; Higgins et al., 2018) (Figure 5b).

386 Sedimentological, geochemical and fluid inclusion data suggest that the majority of
387 primary carbonate minerals precipitating globally were aragonitic in the terminal
388 Ediacaran (Brennan et al., 2004; Cui et al., 2019; Grotzinger et al., 2005; Zhuravlev and
389 Wood, 2008). If aragonite is the dominant carbonate precipitate from the oceans globally,
390 and if the buried carbonate minerals retain a low $\delta^{44}\text{Ca}$ during sediment-buffered
391 recrystallisation, then more ^{40}Ca overall is removed from the ocean, and the $\delta^{44}\text{Ca}$ of the
392 ocean should be higher (Blättler and Higgins, 2017). This increase in $\delta^{44}\text{Ca}_{\text{sw}}$ would mean
393 that the $\delta^{44}\text{Ca}$ in carbonate rocks would be higher on average, as despite the large calcium
394 isotopic fractionation factor, they are precipitating from a fluid with a higher $\delta^{44}\text{Ca}$.
395 Certainly the carbonate minerals would on the whole be higher than the canonical value of
396 -1.5‰ used to signify local primary aragonite deposition in the geological record (Higgins
397 et al., 2018; Husson et al., 2015). It is therefore difficult to explain the very low $\delta^{44}\text{Ca}$

398 captured by some samples from the Nama Group as localized primary aragonite within a
399 calcite-dominated ocean with the same $\delta^{44}\text{Ca}_{\text{sw}}$ as today. However, $\delta^{44}\text{Ca}_{\text{sw}}$ is determined
400 by the final $\delta^{44}\text{Ca}$ of all buried carbonate, which may differ from the primary depositional
401 $\delta^{44}\text{Ca}$, since primary depositional $\delta^{44}\text{Ca}$ may be reset as aragonite recrystallises to calcite.
402 To explain very low $\delta^{44}\text{Ca}$ in ancient carbonate rocks requires *local* aragonite deposition
403 and sediment-buffered $\delta^{44}\text{Ca}$ preservation within oceans where either 1) *globally*, the
404 majority of primary carbonate minerals are deposited as calcite, or 2) *globally*, the majority
405 of carbonate minerals are deposited as aragonite, but recrystallised to calcite or dolomite
406 under fluid-buffered conditions.

407 If we interpret the shift in the $\delta^{44}\text{Ca}$ over the Nama Group as due entirely to local
408 changes in the nature of recrystallization, i.e., in the style of diagenesis, then the shift to
409 lower $\delta^{44}\text{Ca}$ across the Nama Group might record a local change towards recrystallisation
410 under increasingly sediment-buffered conditions (Husson et al., 2015). This could be
411 driven by higher sediment production and accumulation rates, which would push
412 recrystallisation and neomorphism deeper into the sediment pile where it is not in as
413 frequent contact with seawater. Higher burial rates could reflect a higher carbonate
414 saturation state, or an overall marine transgressive succession that created
415 accommodation space to fill with sediment more quickly. Importantly, this interpretation
416 of the shift in $\delta^{44}\text{Ca}$ across the Nama Group would represent a local change in the preserved
417 $\delta^{44}\text{Ca}$ that does not reflect changes in the global ocean, but could provide insight into local
418 depositional conditions and diagenesis in the Nama Group.

419 One way to test whether the trend in $\delta^{44}\text{Ca}$ reflects local changes in fluid- vs.
420 sediment-buffered diagenesis is to compare $\delta^{44}\text{Ca}$ with other geochemical proxies analysed

in the same samples. A transition from fluid-buffered to sediment-buffered diagenetic conditions should produce synchronous changes across multiple geochemical systems (Ahm et al., 2018; Higgins et al., 2018; Husson et al., 2015). For example, in the sulfur isotope system, recrystallisation of carbonate minerals during early diagenesis under fluid-buffered conditions may capture unaltered seawater $\delta^{34}\text{S}$, whereas recrystallisation in sediment-buffered conditions may capture an evolved pore fluid $\delta^{34}\text{S}$ (Rennie and Turchyn, 2014). In the Kuibis Subgroup, there is a weak but significant negative correlation between $\delta^{44}\text{Ca}$ and $\delta^{34}\text{S}_{\text{CAS}}$ ($P < 0.05$, $R^2 = 0.53$; Figure 4k) that supports a change in the realm of diagenesis (sulfur isotope data from Tostevin et al., 2017). However, there is no significant correlation between $\delta^{44}\text{Ca}$ and carbonate-associated-sulfate (CAS) content (p-value = 0.41), which should also be sensitive to fluid- vs. sediment-buffered recrystallisation, since CAS abundance is typically higher in calcite (10,000s of ppm) than in aragonite (1000s of ppm) (Busenberg and Plummer, 1985). Under sediment-buffered conditions, a decrease in CAS abundance would be expected as the sulfate concentration in the fluid would be set by the primary aragonite, but this is not observed.

We can also examine the relationship between $\delta^{44}\text{Ca}$ and uranium isotope data from the Kuibis Subgroup at Zebra River. In modern carbonate sediments, pore water reduction of uranium during early diagenesis offsets $\delta^{238}\text{U}$ in recrystallised carbonate minerals towards higher values (Chen et al., 2018). If recrystallisation takes place under deeper burial conditions, where the supply of uranium is limited, then the bulk carbonate sediment is more likely to retain a $\delta^{238}\text{U}$ close to primary seawater. Changes in the style of diagenesis should hence produce a positive correlation between $\delta^{44}\text{Ca}$ and $\delta^{238}\text{U}$. While in general, the higher $\delta^{44}\text{Ca}$ and $\delta^{238}\text{U}$ both occur in the Lower Omkyk Member, there is no

significant correlation between $\delta^{44}\text{Ca}$ and either $\delta^{238}\text{U}$ (p-value = 0.43) or U/Ca ratios (p-value = 0.17; Figure 4i and 4j) measured on the same samples in the Kuibis Subgroup. However, it is not clear whether this understanding of uranium isotope systematics, developed in modern marine sediments, can be applied to sediments from an ocean with widespread anoxia (Tostevin et al., 2019; Zhang et al., 2018).

During fluid-buffered diagenesis, the Sr content of carbonate rocks should be reduced, reflecting the low abundance of Sr in calcite (1000 ppm). In contrast, sediment-buffered diagenesis can conserve the original high Sr content associated with primary aragonite precipitation (7000–9000 ppm) (Higgins et al., 2018). Changes in the primary mineralogy or style of diagenesis should therefore result in a negative correlation between $\delta^{44}\text{Ca}$ and the Sr content of carbonate rocks (Lau et al., 2017). Overall, there is a negative correlation between strontium content and $\delta^{44}\text{Ca}$ in the Kuibis Subgroup at the $P < 0.05$ level that could support a diagenetic control, but with high scatter ($R^2=0.27$ for a linear trend and $R^2=0.36$ for a log-linear trend). This trend is weak despite efforts to target individual samples with a wide range of strontium contents, including those that are locally anomalous compared to surrounding samples (Figures 2 and 4f). Further, in the Schwarzsrand Subgroup, Sr content in carbonate rocks declines, whereas $\delta^{44}\text{Ca}$ remains low across the section.

In addition, changes in the style of diagenesis can be recorded in oxygen isotope ratio, as $\delta^{18}\text{O}$ in carbonates decreases during recrystallisation at deeper burial depths. Sediment-buffered recrystallisation could therefore result in lower $\delta^{18}\text{O}$ and lower $\delta^{44}\text{Ca}$ (Higgins et al., 2018). We find a weak correlation between $\delta^{44}\text{Ca}$ and $\delta^{18}\text{O}$ at the $P < 0.05$ level ($R^2=0.4$), but this is in the opposite direction than would be expected if these trends

were produced by changes in the style of diagenesis (Higgins et al., 2018; Husson et al., 2015). Such a relationship may be possible if sediment-buffered recrystallisation occurs in the presence of meteoric groundwaters, which can extend at depth below marine continental shelves. However, meteoric cements have not been noted in analysed samples in the Nama Group (Wood et al., 2018). In addition, there is no apparent correlation between $\delta^{44}\text{Ca}$ and $\delta^{13}\text{C}$ or TOC, or with other proxies that are partially impacted by changes in diagenetic conditions, such as Mn content, Mn/Sr, total iron (Fe_T) or Mg/Ca ratios (Figure 4a,e,h).

Times of high fluid flow driving fluid-buffered diagenesis could be expected to occur below sequence boundaries, but the $\delta^{44}\text{Ca}$ trend across the Nama Group cross cuts transgressive systems tracts, high stands and sequence boundaries, similar to $\delta^{44}\text{Ca}$ data in Triassic carbonate rocks (Lau et al., 2017) (Figure 2). This poses a further problem for the widespread interpretation of $\delta^{44}\text{Ca}$ in the geological record to only reflect fluid- vs. sediment-buffered recrystallisation. While local mineralogical and diagenetic controls provide a clear explanation for coupled geochemical trends recorded in modern carbonate sediments (Ahm et al., 2018; Higgins et al., 2018), as well as some ancient carbonate rocks (Ahm et al., 2019; Husson et al., 2015), a diagenetic framework for interpreting $\delta^{44}\text{Ca}$ can only partially explain the geochemical trends across the Nama Group. We suggest that while fluid- vs. sediment-buffered diagenesis is one important way to introduce variability into $\delta^{44}\text{Ca}$ measured in carbonate rocks, it may not be the only driver of change in the $\delta^{44}\text{Ca}$ of ancient carbonate rocks.

5.2 Scenario two: A change in seawater $\delta^{44}\text{Ca}$

490 An alternative suggestion is that the shift in $\delta^{44}\text{Ca}$ seen across the Nama Group
491 represents a change in the global Ca cycle around ~550 Ma that lasted 11–14 Myrs (e.g.,
492 scenario 2; Figure 5c). Assuming there is no systematic change in primary carbonate
493 mineralogy, mineral precipitation rates or diagenetic conditions across the Nama Group,
494 we explore other ways to produce a negative shift in the $\delta^{44}\text{Ca}$ of carbonate rocks. The Late
495 Ediacaran was a time of transformation, including new biological innovations, as well as
496 profound changes in seawater chemistry, climate and style of sedimentation. The marine
497 calcium cycle would likely have been sensitive to each of these changes. We will consider
498 several of these in turn.

499

500 **5.2.1 Evaporite deposition**

501 The removal of calcium from the ocean is divided between carbonate mineral burial,
502 evaporite mineral burial and minor sinks such as alteration of oceanic crust, which can fix
503 calcium into the altered phases. Over long timescales, carbonate minerals are the dominant
504 sink for calcium, and have a $\delta^{44}\text{Ca}$ similar to bulk silicate Earth (Blättler and Higgins, 2017).
505 But during sporadic, geologically short-lived intervals of extensive evaporite deposition,
506 the carbonate sink may form a smaller fraction of the total calcium sink. If enhanced
507 evaporite deposition increases the proportion of calcium buried as CaSO_4 , which typically
508 has a similar $\delta^{44}\text{Ca}$ to seawater due to quantitative removal in evaporite basins (Blättler
509 and Higgins, 2014), then the residual calcium isotope composition of seawater could
510 decrease, through reduced removal of the ^{40}Ca isotope relative to ^{44}Ca . This shift in
511 seawater $\delta^{44}\text{Ca}$ would be reflected in the $\delta^{44}\text{Ca}$ of carbonate minerals, and is consistent
512 with the direction of the $\delta^{44}\text{Ca}$ shift across the Nama Group (Figure 5c).

Mass balance calculations suggest that an increase in the proportional evaporite burial flux (the burial of Ca in evaporites as a proportion of the total Ca burial flux; $F_{\text{evap}}/F_{\text{total}}$), can drive a decrease in seawater $\delta^{44}\text{Ca}$ (Figure 6 and Table 2). A 0.35‰ decrease in seawater $\delta^{44}\text{Ca}$ requires $F_{\text{evap}}/F_{\text{total}}$ to increase from 2.5% to 37%. If the total calcium burial flux remains fixed, at 1.4×10^{13} mols/year, then an $F_{\text{evap}}/F_{\text{total}}$ of 37% equates to an evaporite burial flux of 5.2×10^{12} mols/year. Even extensive evaporite deposition may not generate sufficient calcium fluxes to account for the full magnitude of the $\delta^{44}\text{Ca}$ shift (Hensley, 2006). However, when combined with other changes in the marine Ca cycle such as a change in the style of dolomitisation, a smaller increase in evaporite burial is needed to reconcile the change in $\delta^{44}\text{Ca}$ across the Nama Group.

Extensive evaporite deposition occurred in the late Ediacaran, evidenced by the well-dated sulfate evaporites in Oman and possibly contemporaneous deposits along the northern Gondwana margin in Pakistan, India, Iran and Australia (Claypool et al., 1980; Houghton, 1980; Solomon et al., 1971; Strauss et al., 2001). Radiometric dates and sulfur isotope chemostratigraphy place the deposition of the Kuibis Subgroup in the Nama Group contemporaneous with the A0 Member of the Ara Group, in Oman (Bowring et al., 2007; Fike and Grotzinger, 2008; Tostevin et al., 2017), which contains sulfate evaporite minerals, providing a direct link between the timing of the $\delta^{44}\text{Ca}$ transition and evaporite deposition. Calcite pseudomorphs after gypsum have been reported from the Dengying Formation, South China, which is contemporaneous with the Nama Group (Cui et al., 2019; Duda et al., 2016; Lu et al., 2013).

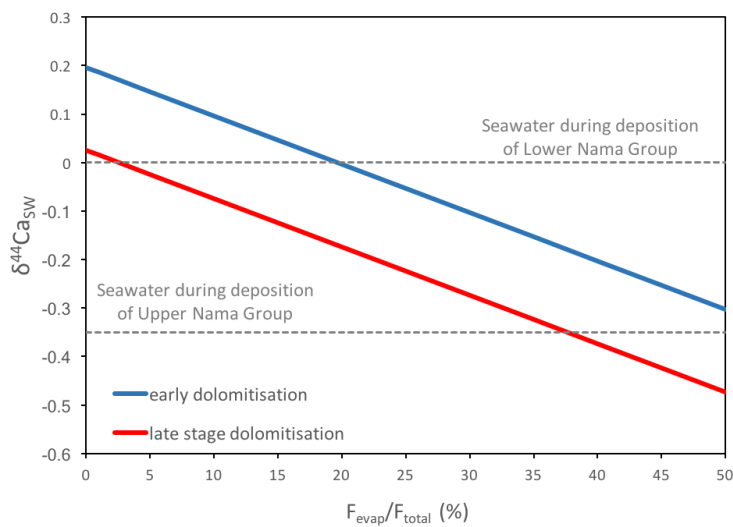


Figure 6: Steady-state mass balance model for seawater $\delta^{44}\text{Ca}$ as a function of the proportional evaporite burial flux ($F_{\text{evap}}/F_{\text{total}}$). Two sets of calculations are shown, using a $\Delta^{44}\text{Ca}$ for dolomitisation of either 0‰ (early dolomitisation; blue line) or -1.58‰ (late stage dolomitisation; red line).

Table 2: Magnitude and $\delta^{44}\text{Ca}$ of fluxes used in evaporite mass balance cycle model.

	Flux	Magnitude	$\delta^{44}\text{Ca}$ (‰)
Inputs	Riverine	1.24×10^{13}	-1.08
	Dolomitisation	1.5×10^{12}	-1.58 (late-stage)
			0 (early)
Outputs	Limestone	$(1 - F_{\text{evap}})$	-1.3 (average)
	Evaporite	F_{evap} (varied from 0 to	-0.18

		100%)	
	Alteration of oceanic crust	$1.5 \cdot 10^{12}$	= seawater

543

544

545 **5.2.2 Enhanced continental weathering**

546 **Enhanced continental weathering could drive an increase in the delivery of calcium**

547 **to seawater.** If there is an imbalance between the riverine Ca flux, and the carbonate

548 sedimentation flux, both the concentration and isotopic ratio of calcium in seawater can be

549 perturbed. For example, a 300% increase in the riverine Ca flux can produce a 0.2–0.4‰

550 negative calcium isotope excursion in seawater over 0.5–1 Myrs (Blättler et al., 2011),

551 which is comparable in magnitude to the progressive 0.35‰ decline in $\delta^{44}\text{Ca}$ across the

552 Nama Group, although over the longer timescale of 11–14 million years. Given the

553 residence time of calcium in seawater (1.1 Myrs in the modern ocean), it is difficult to

554 sustain an isotopic perturbation over such long timescales, because carbonate precipitation

555 rates would rise in response to higher marine calcium concentrations, driving seawater

556 $\delta^{44}\text{Ca}$ back towards higher values.

557 **Enhanced continental weathering is supported by the rise in Sr content across the**

558 **Nama Group. Strontium contents in carbonates are partially controlled by the size of the**

559 **seawater Sr reservoir, which is also sensitive to the mass balance between continental**

560 **weathering and carbonate deposition (Steuber and Veizer, 2002). Although the Sr content**

561 **of carbonate rocks is partially controlled by precipitation rate, mineralogy, and diagenesis,**

562 **global variations in the Sr content of carbonate rocks across multiple sections implies**

secular variation in the size of the marine Sr reservoir. A rise in strontium content through the Kuibis Subgroup is recorded independently in four sections in this study, as well as in previous studies of the Nama Group (Ries et al., 2009), despite the different extraction method used. A similar rise in Sr content been reported globally in the contemporaneous carbonate rocks from the Dengying Formation, south China (Cui et al., 2016b) and the Bambui Formation, Brazil (Caetano-Filho et al., 2019). This supports a secular change in the size of the seawater Sr reservoir consistent with enhanced continental weathering.

5.2.3 Changes in the style of dolomitisation

Over long timescales, the average carbonate depositional sink must be close to bulk silicate Earth, although within this bulk carbonate sink, dolomite tends towards higher $\delta^{44}\text{Ca}$ and limestone towards lower $\delta^{44}\text{Ca}$ (Blättler and Higgins, 2017). Therefore, any changes in the amount of dolomite, or the calcium isotopic fractionation associated with dolomitisation, could influence the $\delta^{44}\text{Ca}$ of seawater, and hence the average $\delta^{44}\text{Ca}$ of limestone deposition. A switch from early dolomitisation, where dolomite retains the low $\delta^{44}\text{Ca}$ associated with primary aragonite or calcite, towards late-stage dolomitisation, where the $\delta^{44}\text{Ca}$ is re-set and ^{40}Ca is preferentially released into pore fluids, could result in a decrease in seawater $\delta^{44}\text{Ca}$ of up to 0.17‰ (Table 2 and Figure 6).

In Neogene environments, dolomite commonly has a higher $\delta^{44}\text{Ca}$ than limestone (Blättler et al., 2015; Fantle and Higgins, 2014; Higgins et al., 2018). In contrast, such differences are not visible in compilations of carbonate rocks across the Precambrian (from 3.0 – 0.7 Ga; Blättler and Higgins, 2017). Early marine cements from carbonate rocks on the Siberian Platform support a fundamental transition in the carbonate system in the Late

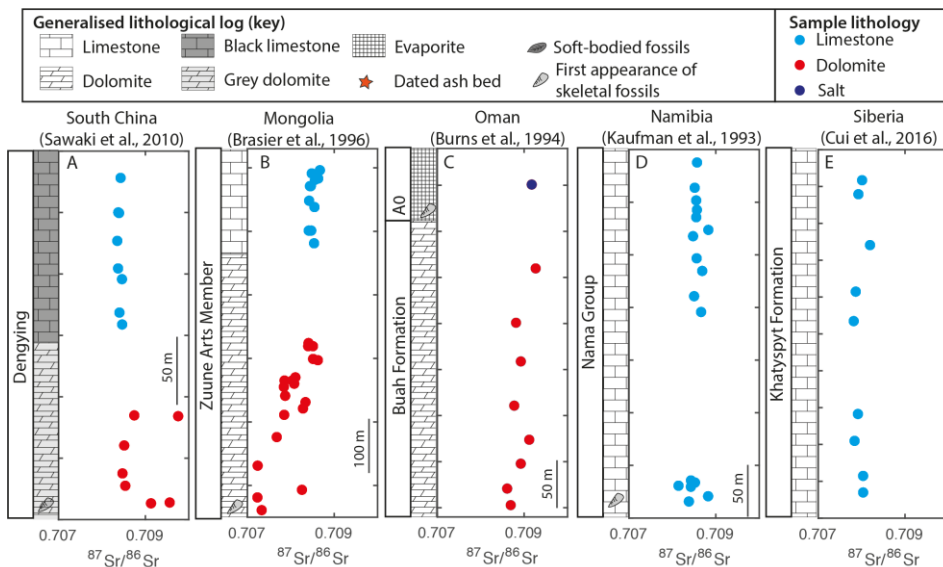
Ediacaran, from “aragonite-dolomite” seas to “aragonite” seas (Wood et al., 2017b), possibly driven by a reduction in the seawater Mg/Ca ratio. It is possible that the late Ediacaran captures a transition in the style of dolomitisation, from early mimetic dolomitisation, with a similar $\delta^{44}\text{Ca}$ to limestone, towards later stage “Phanerozoic style” dolomitisation with a higher $\delta^{44}\text{Ca}$ (Blättler et al., 2015; Fantle and Higgins, 2014). This would drive a decrease in the average $\delta^{44}\text{Ca}$ seawater, which would be reflected in the lower $\delta^{44}\text{Ca}$ captured by marine limestones.

5.2.4 Additional evidence for changes in Ediacaran seawater chemistry

The negative shift in $\delta^{44}\text{Ca}$ across the Nama Group may result from a combination of environmental factors. For example, the rapid change in $\delta^{44}\text{Ca}$ across the Kuibis Subgroup may reflect an imbalance in the calcium cycle during the onset of higher continental weathering rates, resulting in the growth of the marine calcium and sulfate reservoirs, and thus a reduction in the seawater Mg/Ca ratio. The system could then have reached a new steady state in the Schwarzsand Subgroup; with higher weathering rates matched by increased carbonate and evaporite deposition, and changes in the style of dolomitisation. Evaporite deposition could further decrease marine Mg/Ca ratios, as the burial flux of Mg in evaporite deposits, as a proportion of the total Mg budget, is larger than the proportional burial flux of Ca. Increased physical and chemical weathering would impact multiple biogeochemical cycles, delivering Ca, Sr and SO_4^{2-} ions, alkalinity, and nutrients to the oceans (Blättler et al., 2011).

An increase in marine calcium concentrations across the Ediacaran–Cambrian boundary is supported by evidence for a peak in physical and chemical weathering of

continental crust in long-term stratigraphic and geochemical records (Peters and Gaines, 2012). For example, very high $^{87}\text{Sr}/^{86}\text{Sr}$ (from 0.708 to 0.7087) are recorded in Late Ediacaran limestones from South China (Figure 7a), Mongolia (Figure 7b), Oman (Figure 7c), Namibia, (Figure 7d) and Siberia (Figure 7e), interpreted to reflect enhanced continental weathering (Brasier et al., 1996; Burns et al., 1994; Cui et al., 2016a, 2015; Kaufman et al., 1993; Sawaki et al., 2013). Although there is some variability in the $^{87}\text{Sr}/^{86}\text{Sr}$ ratio between sections, the high values are consistent with long term compilations that show $^{87}\text{Sr}/^{86}\text{Sr}$ reached a peak during the late Ediacaran (Halverson et al., 2007). An expansion of seafloor anoxia, recorded by uranium and sulfur isotopes, could be a response to elevated nutrient input (Figure 8) (Tostevin et al., 2019; Zhang et al., 2018). Enhanced delivery of sulfate could also drive a rise in pyrite burial, as well as changes in riverine $\delta^{34}\text{S}$, and could be consistent with $\delta^{34}\text{S}_{\text{CAS}}$ and $\Delta^{33}\text{S}_{\text{CAS}}$ records from the Nama and Ara groups (Figure 8) (Fike and Grotzinger, 2008; Tostevin et al., 2017; Wu et al., 2015). Thus many geochemical proxies have been suggested to reflect some change in terrestrial weathering over this interval.



Commented [RT1]: Petach, works with Emmy smith.
Zuune arts formation

Remove Mongolia
No fossils

Figure 7: $^{87}\text{Sr}/^{86}\text{Sr}$ ratios for the last ca. 10 Myr of the terminal Ediacaran, compiled from the literature. $^{87}\text{Sr}/^{86}\text{Sr}$ from south China are from the Dengying formation (Sawaki et al., 2010). $^{87}\text{Sr}/^{86}\text{Sr}$ from Mongolia were reported from the Tsagaan Oloom Formation (Brasier et al., 1996), and subsequent stratigraphic revisions place the data in the late Ediacaran Zuune Arts Member (Macdonald et al., 2009). Data from Oman are from the Buah and Ara Formations (Burns et al., 1994). Data from the Nama Group are from Kaufman et al., (1993). Data from the Khatyspyt Formation are from (Cui et al., 2016a).

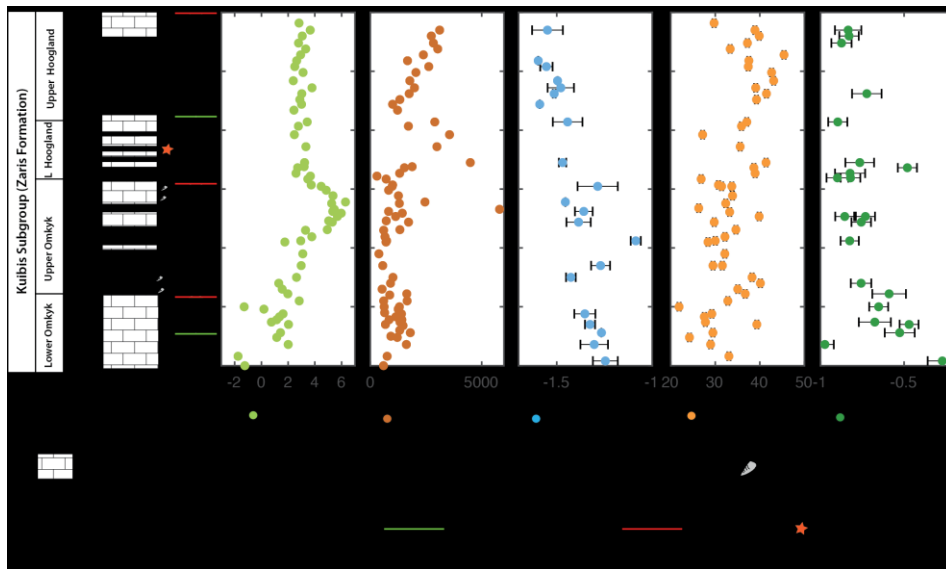


Figure 8: From left to right: Stratigraphic log, sequence stratigraphy, carbon isotope data, Sr content, $\delta^{44}\text{Ca}$, $\delta^{34}\text{S}_{\text{CAS}}$ and $\delta^{238}\text{U}$ for the Lower Nama Group at Zebra River Farm. Beds containing skeletal fossils are marked on the stratigraphic log. $\delta^{238}\text{U}$ and $\delta^{34}\text{S}_{\text{CAS}}$ data are published in full in Tostevin et al., 2019 and 2017, respectively. Sequence stratigraphy is from Adams et al., (2005)

Increased calcium concentrations, and an associated reduction in the seawater Mg/Ca ratio, would increase the carbonate saturation state and alter the stability of various carbonate precipitates. High carbonate saturation states are consistent with a dominance of carbonate lithologies, associated with exceptionally high accumulation rates (~65 m/Myr in the Dengying Formation, and ~100 m/Myr in the Nama Group) (Cui et al., 2016b), as well as densely-aggregating reefs with extensive synsedimentary cement, large meter-scale metazoans, and *Suvorovella* shells beds >1 km in length (Cai et al., 2011;

648 Grotzinger et al., 2005; Penny et al., 2014; Wood et al., 2017a, 2002; Wood and Curtis,
649 2015). This may have been driven by increasing marine calcium concentrations, although
650 the onset of bioturbation around ~560 Ma may also have contributed to increased
651 carbonate saturation, by modifying pH gradients in the top centimetres of sediment and
652 returning sediment alkalinity to bottom waters (Higgins et al., 2009).

653 Despite a proposed increase in sulfate and calcium delivery to the oceans, seawater
654 would likely have remained below critical supersaturation with respect to gypsum.
655 Evaporite deposition is instead controlled by tectonic and climatic factors, such as the
656 formation of semi-isolated basins during rifting. **The assembly of Gondwana, between ~580**
657 **and ~550 Ma, was associated with tectonic inversion and the weathering of Tonian age**
658 **evaporite deposits (Shields et al., 2019). The end of this period may have marked a slow-**
659 **down in evaporite weathering, ushering in a new period of rifting and evaporite deposition.**
660 The rifting of the proposed supercontinent Pannotia **may have occurred around this time,**
661 although the existence of this short-lived supercontinent is debated (Scotese, 2009).
662 Together, geological, sedimentological and geochemical evidence supports higher rates of
663 continental weathering, sustained over tens of millions of years, which delivered sulfate
664 and calcium ions to Late Ediacaran oceans.

665

666 **5.3 Implications for early skeletal animals**

667 Calcareous hard parts appeared relatively abruptly in the terminal Ediacaran (~550
668 Ma) in a range of immobile, shallow marine benthos of probable diverse affinity (Wood,
669 2011). These early skeletal taxa are found exclusively in carbonate settings and likely
670 formed biominerals via a pre-existing organic matrix. Together, these observations suggest

671 that the earliest calcification occurred preferentially in CaCO_3 saturated waters (Wood et
672 al., 2017a). Biomineralisation is energetically costly (Knoll, 2003), and the impetus for its
673 development in the terminal Ediacaran remains enigmatic.

674 If the shift in $\delta^{44}\text{Ca}$ records enhanced continental weathering and delivery of
675 calcium ions, the apparent coincidence in time with the first appearance of skeletal animals
676 may be significant for three reasons. Firstly, high carbonate saturation in shallow shelf
677 environments could have facilitated the onset of passive calcification (Wood et al., 2017a).
678 Secondly, under high seawater calcium concentrations, cellular transporters may struggle
679 to regulate intracellular calcium levels, which can lead to calcium toxicity (Simkiss, 1977).
680 In response, organisms may begin to precipitate carbonate minerals as a mechanism to
681 effectively expel calcium from the cell. Thirdly, a reduction in the Mg/Ca ratio of the ocean
682 would have thermodynamically favoured widespread aragonite precipitation over
683 dolomite (Wood et al., 2017). No organism is known to co-opt dolomite as a biomineral, but
684 most Ediacaran skeletal metazoans formed their shells from aragonite (Zhuravlev and
685 Wood, 2008).

686 While environmental changes may have contributed to the appearance of calcareous
687 skeletons, by making options evolutionarily available that were previously inaccessible, the
688 primary factor explaining the appearance of biomineralisation must be ecological (Vermeij,
689 1989; Wood, 2011). Biomineralisation was preceded by the appearance of motility (~560
690 Ma) and possible carnivory (~550 Ma), and the development of hard parts would have
691 been beneficial for the protection of soft tissue (Knoll, 2003). However, it is possible that
692 environmental changes made building aragonitic skeletons progressively more favourable
693 after ~550 Ma.

694

695 **6. Conclusions**

696 We present new $\delta^{44}\text{Ca}$ data for limestone rocks from the Nama Group, Namibia, that
697 reveal a shift towards lower $\delta^{44}\text{Ca}$ around ~550 Ma. One interpretation of this shift is a
698 local change from fluid- to sediment-buffered diagenesis of primary aragonite deposited
699 over this interval. However, this interpretation is difficult to reconcile with changes in
700 other geochemical and sequence stratigraphic records from the Nama Group that change
701 across independent timescales. If instead, $\delta^{44}\text{Ca}$ records changes in the marine calcium
702 cycle, the data could indicate enhanced weathering fluxes, matched by increased evaporite
703 deposition and changes in the style of carbonate deposition. Enhanced weathering would
704 also deliver more SO_4^{2-} , alkalinity and nutrients to the oceans, and this is supported in the
705 late Ediacaran by an array of stratigraphic, sedimentological and geochemical records.
706 Increased marine calcium concentrations, and lower Mg/Ca ratios (<5) could have
707 facilitated the appearance of aragonitic skeletal animals, which are documented from
708 within the same successions.

709

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715

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